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NEWS	18	JAN	26	Updated MeSH vocabulary, new structured abstracts, and other enhancements improve searching in STN reload of MEDLINE
NEWS	19	JAN	28	CABA will be updated weekly
NEWS	20	FEB		PCTFULL file on STN completely reloaded
NEWS	21	FEB		STN AnaVist Test Projects Now Available for Qualified Customers
NEWS	22	FEB	25	LPCI will be replaced by LDPCI
NEWS	EXP			DECEMBER 2010 CURRENT WINDOWS VERSION IS V8.4.2 .1, WENT DISCOVER FILE IS DATED 24 JANUARY 2011.
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=> d 11

- ANSWER 1 OF 1 REGISTRY COPYRIGHT 2011 ACS on STN
- 11121-48-5 REGISTRY RN
- ED Entered STN: 16 Nov 1984
- Rose Bengal (CA INDEX NAME) CN
- OTHER NAMES:
- CN Bengal Rose CN
- Rose Bengale
- MF Unspecified
- CT COM, MAN
 - STN Files: ADISINSIGHT, AGRICOLA, BIOSIS, BIOTECHNO, CA, CABA, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CIN, DDFU, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IMSRESEARCH, IPA, MEDLINE, MSDS-OHS, PIRA, REAXYSFILE*, RTECS*, TOXCENTER, USPAT2, USPATFULL, USPATOLD

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*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3065 REFERENCES IN FILE CA (1907 TO DATE)
123 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
3080 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> file caplus COST IN U.S. DOLLARS

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FILE COVERS 1907 - 3 Mar 2011 VOL 154 ISS 10 FILE LAST UPDATED: 2 Mar 2011 (2010302/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Oct 2010 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Oct 2010

CAplus now includes complete International Patent Classification (IPC) reclassification data for the fourth quarter of 2010.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

(TUMOR OR TUMORS) 5315 TUMOUR 1968 TUMOURS 7151 TUMOUR

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L7 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2011 ACS on STN
ACCESSION NUMBER:
                       2004:220149 CAPLUS
DOCUMENT NUMBER:
                        140:266883
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L3

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L6

TITLE: Chemotherapy method using x-rays
INVENTOR(S): Wang, Chia-gee; Helson, Lawrence
PATENT ASSIGNEE(S): Nanodaynamics, Inc., USA
SOURCE: PCT Int. Appl., 41 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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AB A method of treating cancer in a human uses x-

rays to disrupt a linkage in a complex of a chemotherapeutic agent and a carrier compound comprising a preselected element. The complex is administered to the human and then a localized region of cells which contains the cancerous cells is irradiated with line emission x-rays of an energy selected to cause emission of Auger

electrons from the pre-selected element of the carrier compound to disrupt the linkage and release the chemotherapeutic agent near the cancer ${}^{\prime}$

cells. A kit useful for the treatment comprises an x-ray tube capable of emitting monochromatic line emission

x-rays and the complex compound A transfer compound useful

in the method comprises a chemotherapeutic agent linked to a carrier compound

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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FILE 'REGISTRY' ENTERED AT 12:30:46 ON 03 MAR 2011

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FILE 'CAPLUS' ENTERED AT 12:31:06 ON 03 MAR 2011
L2 3080 S L1
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L3 90 S L2 AND (CANCER OR TUMOR OR TUMOUR OR NEOPLASM)

L4 22 S L3 AND (RADIATION OR RADIOTHERAPY OR X-RAY OR IRRADIATION)

L5 22 DUP REM L4 (0 DUPLICATES REMOVED)

L6 22 S L5

L7 1 S L5 AND (MONOCHROMATIC OR AUGER)

L6 ANSWER 1 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2010:382585 CAPLUS

DOCUMENT NUMBER: 152:373811

TITLE: Intracorporeal medicaments for high energy

phototherapeutic treatment of disease
INVENTOR(S): Dees, H. Craig; Scott, Timothy C.; Wachter, Eric A.;

Fisher, Walter G.; Smolik, John

PATENT ASSIGNEE(S): Provectus Pharmatech, Inc., USA

SOURCE: U.S. Pat. Appl. Publ., 15pp., Cont.-in-part of U.S.

Ser. No. 542,533. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 13 PATENT INFORMATION:

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US	6331	286			B1						1998-					9981	221
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AI	35/9	100			т.		2007			HT.	2001-	0266	02		2	0010	403
ES	2005	0207	076		7.1		2007			HC.	2001- 2005- 2006-	1246	U Z E /I		2	0010	600
110	2003	0207	076		7.1		2003	0722		TTC	2005-	5/25	33		2	0030	002
PRIORIT					AI		2007	0403		HS.	1998-	2167	87		12 1	9981	221
11/10//11	1 111 1	DI4.	1141 0	• •						IIS	2000-	1950	900		2	0000	406
										IIS	2000- 2001-	8174	48		A2 2	0010	326
										US	2006-	5425	3.3			0061	
										US	1996-	7413	70			9961	
										WO	1997-	US19:	249	1	W 1	9971	027
										US	1997-1 1998-	9683	2		A 1	9980	612
							2007			WO	1999-	US12	056	1	7 I	9990	528
										US	1999-	3826	22		A3 1	9990	825
										US	2000-	1879	58P	1	P 2	0000	309
										US	2000- 2001- 2001-	7798	8 0		A 2	0010	208
										WO	2001-	US72	31	1	vi 2	0010	307

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 152:373811

AB New intracorporeal radiodense medicaments and certain medical uses and

methods for use of such high energy phototherapeutic medicaments for treatment of human or animal tissue are described, wherein a primary active component of such medicaments is a halogenated xanthene or halogenated xanthene derivative The halogenated xanthenes constitute a family of potent radiosensitizers that become photoactivated upon irradiation of the treatment site with ionizing radiation. In embodiments of the present invention, such medicaments are used for treatment of a variety of conditions affecting the skin and related organs, the mouth and digestive tract and related organs, the urinary and reproductive tracts and related organs, the respiratory tract and related organs, the circulatory system and related organs, the head and neck, the endocrine and lymphoreticular systems and related organs, various other tissues, such as connective tissues and various tissue surfaces exposed during surgery, as well as various tissues exhibiting microbial or parasitic infection. In another embodiment, such medicaments are produced in various formulations including liquid, semisolid, solid or aerosol delivery vehicles.

THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD

5 (5 CITINGS) ANSWER 2 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2010:175815 CAPLUS

DOCUMENT NUMBER: 152:247629

TITLE: Composition for a tissue repair implant and methods of making the same

INVENTOR(S): Chen, Silvia S.; Chen, Jingsong; Wolfinbarger, Lloyd, Jr.

PATENT ASSIGNEE(S):

OS.CITING REF COUNT:

SOURCE: U.S. Pat. Appl. Publ., 24pp.

CODEN: USXXCO DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

	TENT				KIN	D	DATE			APPL						ATE	
						_											
US	2010	0036	503		A1		2010	0211		US 2	008-	1881	27		2	0080	807
WO	2010	0169	42		A1		2010	0211		WO 2	009-	US45.	56		2	0090	807
	W:	ΑE,	AG,	AL,	AM,	AO,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,
		CA,	CH,	CL,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,
		ES,	FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,
	ES, FI, KE, KG, MD, ME, PG, PH, SY, TJ,				KN,	KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,
		MD,	ME,	MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PE,
		PL,	PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	ST,	SV,		
		TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW		
	PG, PH, SY, TJ, RW: AT, BE,					CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HR,	HU,
		IE,	IS,	IT,	LT,	LU,	LV,	MC,	MK,	MT,	NL,	NO,	PL,	PT,	RO,	SE,	SI,
		SK,	SM,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,
		SN,	TD,	TG,	BW,	GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	ŞD,	SL,	SZ,	TZ,	UG,
		ZM,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM					
US	2011		A1		2011	0224		US 2	010-	7329	74		2	0100	326		
PRIORIT	Y APP	INFO	. :						US 2	005-	2472	30	- 1	A1 2	0051	012	
										US 2	008-	1881	27	- 2	A 2	0080	807
										US 2	009-	3946	29		A2 2	0090	227

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

The invention is directed to a process for making a tissue repair implant having a porous sponge-like structure to repair bone, cartilage, or soft tissue defects. A process for preparing a biol. functional tissue repair implant comprises steps of (a) producing a connective tissue homogenate from one or more connective tissues, (b) mixing the connective tissue homogenate with a carrier solution to produce a connective tissue carrier, (c) optionally mixing one or more natural or synthetic bone fragments with said connective tissue carrier to produce a tissue repair mixture, (d) freezing or freeze-drying the tissue repair mixture to produce a porous sponge-like structure and create a three-dimensional framework to entrap the natural or synthetic bone fragments, and (e) treating the frozen or freeze-dried porous sponge-like structure with one or more treatment solns. to produce a stabilized porous sponge-like structure. A crudely fragmented connective tissue from one or more connective tissues is optionally mixed with the tissue repair mixture before freezing or freeze-drving. The tissue repair implant having a porous sponge-like structure is optionally combined with one or more bioactive supplements or one or more agents that have bioactive supplement binding site(s) to increase the affinity of growth factors, differentiation factor, cytokines, or anti-inflammatory agents to the tissue repair implant. The invention is further directed toward applying such tissue repair implant for tissue repair. Thus, homogenized fascia lata was mixed with a sodium alginate solution to produce a connective tissue carrier that was mixed further with crudely fragmented fascia and sized, freeze-dried demineralized bone matrix (DMB) powder. The mixture was distributed into molds with predetd. shapes and sizes, freeze-dried, treated with CaCl2, washed with water, freeze-dried again, optionally exposed to a neg. hydrostatic pressure to allow the expansion of the DMB mixture to a preset thickness, and sterilized. The freeze-dried, molded, tissue repair implants obtained were porous sponge-like structure with DMB particles having high mech. strength and maintaining the shape of their mold.

L6 ANSWER 3 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2010:175814 CAPLUS

DOCUMENT NUMBER: 2010:175814 CA

TITLE: Composition for a tissue repair implant and methods of making the same

INVENTOR(S): Chen, Jingsong; Wolfinbarger, Lloyd; Chen, Silvia S. PATENT ASSIGNEE(S): LifeNet Health, USA

PATENT ASSIGNEE(S): LifeNet Health, USA SOURCE: PCT Int. Appl., 70 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

	TENT :				KIN	D	DATE			APPL	ICAT	ION I	. 01/		D.	ATE	
						_									-		
WO	2010	0169	42		A1		2010	0211		WO 2	009-	US45.	56		2	0090	807
	W:	ΑE,	AG,	AL,	AM,	AO,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,
		CA,	CH,	CL,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,
		ES,	FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,
		KE,	KG,	KM,	KN,	KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,
		MD,	ME,	MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PE,
		PG,	PH,	PL,	PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	ST,	sv,
		SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HR,	HU,
		ΙE,	IS,	IT,	LT,	LU,	LV,	MC,	MK,	MT,	NL,	NO,	PL,	PT,	RO,	SE,	SI,
		SK,	SM,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,
		SN,	TD,	TG,	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,
		ZM,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM					
US	2010	0036	503		A1		2010	0211		US 2	-800	1881	27		2	0800	807
RIT	Y APP	LN.	INFO	. :						US 2	008-	1881	27		A 2	0080	807

PRIORITY APPLN. INFO.: US 2008-188127 A ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

The invention is directed to a process for making a tissue repair implant having a porous sponge-like structure to repair bone, cartilage, or soft tissue defects. A process for preparing a biol. functional tissue repair implant comprises steps of (a) producing a connective tissue homogenate from one or more connective tissues, (b) mixing the connective tissue

homogenate with a carrier solution to produce a connective tissue carrier, (c) optionally mixing one or more natural or synthetic bone fragments with said connective tissue carrier to produce a tissue repair mixture, (d) freezing or freeze-drying the tissue repair mixture to produce a porous sponge-like structure and create a three-dimensional framework to entrap the natural or synthetic bone fragments, and (e) treating the frozen or freeze-dried porous sponge-like structure with one or more treatment solns, to produce a stabilized porous sponge-like structure. A crudely fragmented connective tissue from one or more connective tissues is optionally mixed with the tissue repair mixture before freezing or freeze-drying. Thus, homogenized fascia lata was mixed with a sodium alginate solution to produce a connective tissue carrier that was mixed further with crudely fragmented fascia and sized, freeze-dried demineralized bone (DMB) powder. The mixture was distributed into molds with predetd. shapes and sizes, freeze-dried, treated with CaCl2, washed with water, freeze-dried again, optionally exposed to a neg. hydrostatic pressure to allow the expansion of the DMB mixture to a preset thickness, and sterilized. The freeze-dried, molded, tissue repair implants obtained were porous sponge-like structure with DMB particles having high mech. strength and maintaining the shape of their mold.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2009:918420 CAPLUS DOCUMENT NUMBER: 151:205597

TITLE:

Wearable photoactivator for ocular therapeutic applications and uses thereof for treatment of ocular

disease including infection, neoplasia, and corneal dystrophies

INVENTOR(S): Soltz, Robert; Soltz, Barbara Ann; Behrens, Ashley

PATENT ASSIGNEE(S): The Johns Hopkins University, USA

SOURCE: U.S. Pat. Appl. Publ., 24pp. CODEN: USXXCO

Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION: PATENT NO.

PATENT NO.		DATE	APPLICATION NO.	
US 20090192437	A1	20090730		20080924
PRIORITY APPLN. INF	0.:		US 2007-994979P	P 20070924
ASSIGNMENT HISTORY	FOR US PATEN	T AVAILABLE	E IN LSUS DISPLAY FOR	MAT
AB The invention	provides a w	earable de	vice for delivery of	light of a
			cornea of a subject.	
includes a fra	me for attac	hment of a	light source housing	which includes a
			n the housing to allo	
			and the light source	
			tion provides method	
			isease including infe	
			of the invention can	
			utic agents. Thus, p	
			as fitted with a wear	
			ng a UV-A light sourc	
			ide light over 3 to 1	
			exposed, based on th	
			is such that it warra	
			ea before penetrating	
			the exposure of other	
			gh an opening in the	
riboflavin to	the eye in t	he form of	drops and the ribofl	avin solution

is in the range of about 0.1 % to 5 % to completely bathe the eye in riboflavin.

L6 ANSWER 5 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2008:914722 CAPLUS

DOCUMENT NUMBER: 151:191670

TITLE: Comparison on photodynamic actions of AIPcS2 and Rose

Bengal on erythrocytes

AUTHOR(S): Zhorina, L. V.; Zmievskii, G. N.

CORPORATE SOURCE: N. E. Bauman Moscow State Technical University, Moscow, Russia

SOURCE: Tekhnologii Zhivykh Sistem (2008), 5(2-3), 51-56

CODEN: TZSEAC

PUBLISHER: Izdatel'stvo "Radiotekhnika"

DOCUMENT TYPE: Journal LANGUAGE: Russian

The search for new photosensitizes (PS) for traditional purposes and new fields of photodynamic action (PDA) is being carried out now. At the same time the effectiveness of different Pc action in similar conditions is compared. Rose Bengal (RB) is known as Pc with high quantum output of singlet oxygen ($\phi = 0.76$) and is characterized by a set of destroying mechanisms in case the PDA. Deficiency of RB is absorbing maximum at green field of spectra (520 and 560 nm). Nevertheless RB is effective PS for different tissues (including cancer) and for red blood cells. Sulfonated aluminum phthalocyanine has more suitable for PDA intensive absorb maximum in far red field of spectra (670...680 nm), high quantum yield of singlet oxygen (up to 0,5), high accumulation level in tumor tissues in comparison with normal ones, is removing from organism quite rapidly. The comparison of the photodynamic action on erythrocytes AlPcS2 and Rose Bengal is presented. The following events are possible at PDA: erythrocytes geometry changing, breaking of membrane and erythrocyte's destruction. At the same time erythrocytes are prevailing among others blood elements therefore they determine optical, mech. and other properties of blood. So, radical changes of optical blood properties (absorption, scattering) should be expected. The optical transparency of erythrocyte suspension at PDA was measured. It was discovered that (1) erythrocytes with accumulated PS die at low irradiation doses; (2) erythrocytes incubated and nonincubated with RB die at higher irradiation doses than with AlPcS2 ones. Point out that absorb maximum of oxyHb are at 540 and 576 nm, so they are very close to absorb maximum of RB. This "neighborhood" may lead to catching the source radiation energy by Hb instead of RB. Probably this is the reason of the second result of our investigation. External appearance of erythrocytes was under visual control. It was revealed that at in rise transparency the erythrocytes form at first became spherical, then it looked like a volume star, after that erythrocytes were destroyed and disappeared from the field of vision of the microscope. So we conclude that the form changes and the following gemolyze of erythrocytes have place because of osmotic pressure changes due to the destruction of membrane transport, breaking barrier properties and permeability of membrane. The fact that AIPcS2 causes photodynamic effect at much less doses of irradiation than Rose Bengal is shown. If our idea about catching the source radiation energy by Hb instead of RB

L6 ANSWER 6 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2006:976194 CAPLUS

ACCESSION NUMBER: 2006:976194 DOCUMENT NUMBER: 145:328416

TITLE: Ellagic acid-related compound and polyaromatic phenol

is correct, we can say that the use of RB as PS for PDA is not effective.

inhibitors of glutathione-S-transferase, and their

therapeutic use

INVENTOR(S): Becker-Brandenburg, Katja; Zimmermann, Herbert;

Fritz-Wolf, Karin

PATENT ASSIGNEE(S): Universitaet Giessen, Germany; Max-Planck-Gesellschaft Zur Foerderung der Wissenschaften e.v.

PCT Int. Appl., 66pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. WO 2006097472 A2 20060921 WO 2006-EP60707
WO 2006097472 A3 20070907 ---- ------ ------- ------W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, AI, BE, BG, CH, CI, CZ, DE, DR, EE, ES, FI, FR, GB, GR, RU, E, CIS, TI, TL, LU, LV, MC, ML, PI, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, GRG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA

KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA

EP 1865942 A2 20071219 EP 2006-708757 20060314

R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, MK, YU A 20071109 IN 2007-DN7684 IN 2007DN07684 20071008 IN 2007-DN7684 20071008 US 2005-661596P P 20050314 WO 2006-EP60707 W 20060314 PRIORITY APPLN. INFO.:

OTHER SOURCE(S): MARPAT 145:328416

AB The invention discloses methods for preventing, treating, or ameliorating medical conditions, including cancer, drug resistance, and parasite infections such as malaria, by administering compds. that are capable of inhibiting glutathione-S-transferase (GST), as well as to the use of these compds. for preparing pharmaceutical compns. for preventing, treating, or ameliorating the medical conditions. Furthermore, the invention discloses ellagic acid-related compound and polyarom, phenol inhibitors of GST, as well as pharmaceutical compns. comprising these GST inhibitors, optionally comprising further compds. known to be effective in treating the medical conditions.

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L6 ANSWER 7 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2005:17009 CAPLUS

142:107447 DOCUMENT NUMBER:

TITLE: Bivalent inhibitors of glutathione transferases

Lyon, Robert P.; Atkins, William M.; Maeda, Dean Y.; INVENTOR(S):

Zebala, John A. PATENT ASSIGNEE(S):

USA U.S. Pat. Appl. Publ., 33 pp. SOURCE:

CODEN: USXXCO DOCUMENT TYPE: Pat.ent.

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

A1 20050106 US 2004-878732 20040628 .: US 2003-483320P P 20030627 US 20050004038 PRIORITY APPLN. INFO.:

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 142:107447

AB Bivalent inhibitors having affinity for one or more dimeric glutathione-S-transferase (GST) isoenzymes are provided. The bivalent inhibitors comprise two ligand domains connected by a mol. linker, wherein the ligand domains have affinity for one or more monomers in the one or more dimeric GST isoenzymes. The ligand domains are separated by a distance ranging from about 5 to about 100 Å. The bivalent inhibitors of the invention demonstrate greatly improved affinity for GST isoenzymes. In a specific embodiment, the bivalent inhibitors of the invention further provide affinity for substantially one GST isoenzyme and for substantially one GST class. The bivalent inhibitors of the invention have numerous uses that include the treatment of drug-resistant cancer, malaria, and stimulation of hematopoiesis. For example, an IC50 was determined for each of the C16-20 bis(glutathionyl)alkyl esters (preparation given) with GST isoenzymes Al-1 and Pl-1. An IC50 was also determined for the monovalent inhibitor. Notably, each of the bis(qlutathionyl alkyl)esters exhibited an IC50 more than one order of magnitude lower than the monovalent benchmark compound and six orders of magnitude lower than Km of glutathione. From this data, it is evident that the bivalent inhibitors exhibit between 10- and 100-fold greater affinities than the corresponding monovalent inhibitor. Different affinities of the bivalent inhibitors for the

GSTP1-1 and GSTA1-1 isoenzymes were observed 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD OS.CITING REF COUNT: (3 CITINGS)

L6 ANSWER 8 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2004:220149 CAPLUS

DOCUMENT NUMBER: 140:266883

TITLE: Chemotherapy method using x-rays INVENTOR(S): Wang, Chia-gee; Helson, Lawrence

PATENT ASSIGNEE(S): Nanodaynamics, Inc., USA SOURCE: PCT Int. Appl., 41 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent.

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PA:	TENT :	NO.			KIN	D	DATE			APPL	ICAT	ION I	NO.		D	ATE	
		2004	0219	82		A2 A3		2004			WO 2	003-	US27.	242		2	0030	903
		W:	co,	CR,	CU,	CZ,	DE,	AU, DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
						LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,
		PG, PH, PL TR, TT, TZ RW: GH, GM, KE			TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
		RW:	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		0001	BF,	ВJ,		CG,	CI,	IE, CM,	GA,	GN,	GQ,	GW,	ML,	MR,		SN,	TD,	TG
	US 20040259811 AU 2003278748					A1 A1		2004 2004			AU 2	003-	2787	48		2	0030	903
PRIOR	RIORITY APPLN. INFO.:										US 2	003-	6513	07		A 2	0020	828
											WO 2	UU.3-	1527	242		N 2	0030	20.5

AB A method of treating cancer in a human uses xrays to disrupt a linkage in a complex of a chemotherapeutic agent

and a carrier compound comprising a preselected element. The complex is administered to the human and then a localized region of cells which contains the cancerous cells is irradiated with line emission xrays of an energy selected to cause emission of Auger electrons from the pre-selected element of the carrier compound to disrupt the linkage and release the chemotherapeutic agent near the cancer cells. A kit useful for the treatment comprises an x-ray tube capable of emitting monochromatic line emission x-rays and the complex compound A transfer compound useful in the method comprises a

chemotherapeutic agent linked to a carrier compound 2 REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2002:964924 CAPLUS

DOCUMENT NUMBER: 138:44708

TITLE. Polymer gel for cancer treatment

Zheng, Ji; Chu, Feng INVENTOR(S):

USA PATENT ASSIGNEE(S):

SOURCE: U.S. Pat. Appl. Publ., 13 pp.

CODEN: USXXCO DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. ----------US 20020192289 A1 20021219 US 2002-173354 20020615 PRIORITY APPLN. INFO:: US 2001-298943P P 20010618

AB A method is disclosed for cancer treatment based on using a

solid polymer gel to completely block blood vessels of tumor. A

polymer aqueous solution is injected into blood vessels and formed a solid gel

in blood vessels of tumor by applying electromagnetic radiation or temperature source at tumor tissue to inducing

crosslinking or phase transition. The tumor cells starve and perish because of without nutrients and oxygen provided by vascularization

and metastasis can also be prevented because polymer gels blocks

tumor cells to shed into blood circulation, when the blood vessels of tumor are completely blocked by the solid polymer gels.

Also, anti-cancer drug including chemotherapy drug,

radiation drug or anti-angiogenic drug can be mixed or conjugated

with the polymer in polymer aqueous solution to be locally delivered to the tumor after polymer gel formation in the blood vessels of

tumor of human or animal. An example photopolymerizable polymer is branched PEG-cinnamylideneacetyl chloride.

L6 ANSWER 10 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2002:240566 CAPLUS DOCUMENT NUMBER: 136:241657

Phototherapeutic and chemotherapeutic immunotherapy TITLE:

against tumors
Dees, H. Craig; Scott, Timothy; Wachter, Eric
Photogen, Inc., USA
PCT Int. Appl., 23 pp. INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

CODEN: PIXXD2 Patent

DOCUMENT TYPE:

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO.
                KIND DATE APPLICATION NO. DATE
    WO 2002024199
                      A1 20020328 WO 2001-US29179 20010919
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
            ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    US 20020107281 A1 20020808
                                       US 2001-952448
    AU 2001096258
                      A
                            20020402
                                        AU 2001-96258
                                                             20010919
PRIORITY APPLN. INFO.:
                                        US 2000-234654P
                                                         P 20000922
                                        US 2001-952448
                                                         A 20010914
                                        WO 2001-US29179
                                                         W 20010919
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

AB The present invention is directed to new methods, medicaments and
pharmaceutical compns. for improved cancer treatment that lower
recurrence of the primary tumor by causing selective, acute
destruction of tumor tissue and thereby exposing the immune
system to large amts. of substantially non-denatured tumor
material over a short period of time. Several examples are provided in
which phototherapy, Rose Bengal, or a combination of Rose Bengal and
radio-/phototherapy were used in animals to enhance the body's immune
system to elicit an antitumor immune response.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 11 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2001:416760 CAPLUS

DOCUMENT NUMBER: 135:16142

TITLE: Radiation-absorbing dyes for treating

illnesses associated with abnormal vasculature

INVENTOR(S): Flower, Robert W.; Alam, Abu

PATENT ASSIGNEE(S): Akorn, Inc., USA SOURCE: PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

		ENT:				KIN	D	DATE		- 1	APPL:	ICAT:	ION	NO.		D	ATE		
	WO	2001 2001	0397	64		A2 A3		2001		1	WO 2	000-	US41	110		2	0001	010	
	WO		AE, CR, HU, LU,	AG, CU, ID, LV, SE,	AL, CZ, IL, MA,	AM, DE, IN, MD,	AT, DK, IS, MG,	AU, DM, JP, MK, SL,	AZ, DZ, KE, MN,	EE, KG, MW,	ES, KP, MX,	FI, KR, MZ,	GB, KZ, NO,	GD, LC, NZ,	GE, LK, PL,	GH, LR, PT,	GM, LS, RO,	HR, LT, RU,	
		RW:	AT,		CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	
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conjugates of any of the foregoing dyes) for the treatment of conditions associated with abnormal vasculature, including lesions, and, more specifically, tumors (cancerous and benign) and choroidal neovascularization (CNV) associated with age-related macular degeneration (ARMD) is described. A method for treating a lesion in an animal having a blood vessel that carries blood into the lesion, comprises administering a first composition containing the above photodynamic dye, and a carrier to fill

at

least a portion of the lesion with the first composition Radiation is applied to the photodynamic dye in the lesion of a type and in an amount sufficient to excite the photodynamic dye, and applying radiation to the blood vessel in an amount sufficient to increase the temperature of the vessel.

REFERENCE COUNT:

4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 12 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

Patent.

ACCESSION NUMBER: 2000:513548 CAPLUS

DOCUMENT NUMBER: 133:131883

TITLE: Method for improved radiation therapy

INVENTOR(S): Wachter, Eric; Smolik, John; Dees, H. Craig

PATENT ASSIGNEE(S): Photogen, Inc., USA SOURCE: PCT Int. Appl., 27 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PF	TENT	NO.			KIN	D	DATE			APPI	ICAT	ION I	NO.		D	ATE	
WC	2000	0430	45		A1		2000	0727		WO 2	000-	JS18	15		2	0000	125
	W:	ΑE,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
		CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
		IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,
		MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,
		SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,	ZA,	ZW			
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,	CF,
		CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG				
CF	2358	989			A1		2000	0727		CA 2	000-	2358	989		2	0000	125
EF	1146	912			A1		2001	1024		EP 2	-000	9083	66		2	0000	125
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO										
BF	2000	0076	92		A		2001	1106		BR 2	-000	7692			2	0000	125
JE	2002	5352	91		T		2002	1022		JP 2	-000	5944	98		2	0000	125
/I	2001		A		2005	0304		IN 2	001-	CN10	07		2	0010	717		
IN	2001	CN01	807		A		2005	0520		IN 2	001-	CN18	07		2	0010	717
M	2001	0074	87		A		2001	1203		MX 2	001-	7487			2	0010	725
PRIORIT	Y APP	LN.	INFO	. :						US 1	999-	2362	47	- 2	A 1	9990	125
										WO 2	000-	JS18	15	1	W 2	0000	125

- AB A method is disclosed for treating a selected volume of tissue which method includes distributing a radiosensitizer and a plurality of ionizing radiation sources substantially within the volume of tissue to produce treatment zones that are generally uniformly distributed throughout the volume of tissue. An agent is also disclosed for treating such tissue, wherein the agent includes a radiosensitizer and an ionizing radiation source used in conjunction to define an injectable
- treatment agent.

 OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)
- REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

L6 ANSWER 13 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1997:156828 CAPLUS DOCUMENT NUMBER: 126:235320

ORIGINAL REFERENCE NO.: 126:45472h, 45473a

TITLE: Comparative studies on the tolerance to photoinduced

cutaneous inflammatory reactions by psoralen and rose

AUTHOR(S): Kumar, Janak R.; Haberman, Herbert F.; Ranadive,

Narendranath S.

CORPORATE SOURCE: Department of Medicine, University of Toronto,

Toronto, ON, M5S 1A8, Can.

SOURCE: Journal of Photochemistry and Photobiology, B: Biology

(1997), 37(3), 245-253 CODEN: JPPBEG; ISSN: 1011-1344

Elsevier PUBLISHER . DOCUMENT TYPE: Journal

LANGUAGE: English

The photochemotherapeutic value of topical 8-methoxypsoralen (8-MOP) plus UVA irradiation has been well recognized. The phototoxicity associated with psoralen plus UVA (PUVA) therapy is hallmarked by an increase in vascular permeability (iVP), the accumulation of polymorphonuclear leukocytes (aPMN) and erythema formation in situ. Rose bengal (RB) plus UVA-VIS light (320-700 nm) produces a similar acute inflammatory response, but without immediate or delayed erythema and perceptible edema. This study describes some of the parameters involved in inflammatory reactions evoked by PUVA and the results are compared with RB-induced phototoxic reactions. The rates of iVP and aPMN with a 3 h pulse were quantified using 125I-albumin and 51Cr-labeled PMNs resp. The erythemal response was graded visually. 8-MOP cream was applied topically, while RB was injected intradermally in rabbit skin before UVA-VIS (9.4 J cm-2) irradiation The data show that there is no significant difference in the rates of iVP, aPMN and erythema formation between normal skin sites and mast cell-depleted skin sites when challenged with 8-MOP plus light. These results suggest that in situ mast cells do not play a significant role in 8-MOP-photoinduced acute cutaneous inflammatory reactions, in contrast with RB-photoinduced reactions. The iVP and aPMN responses are minimal or absent in sites subjected to repeated exposure to 8-MOP plus light for three or more consecutive days, suggesting the establishment of a desensitized/unresponsive state. Moreover, 8-MOP-photo-desensitized sites do not produce iVP and aPMN of the same magnitude as the normal (naive) skin sites when challenged with RB plus light. Similarly, RB-photo-desensitized sites do not produce iVP and aPMN of the same magnitude as the native skin sites when challenged with 8-MOP plus light. The desensitization and cross-desensitization of skin sites to 8-MOP- or RB-photoinduced reactions suggest that there is either direct attack on the target cell(s), thereby removing the ability to express adhesion mols., such as endothelial leukocyte adhesion mol. 1 (ELAM-1) or intercellular adhesion mol. 1 (ICAM-1), involved in the accumulation of inflammatory cells, or downregulation of the secretion/release of putative agent(s), such as interleukin 1 (IL-1) and tumor necrosis factor α (TNF- α), responsible for the initiation and progression of cutaneous inflammations.

OS.CITING REF COUNT: THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

ANSWER 14 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1997:43829 CAPLUS DOCUMENT NUMBER: 126:154514

ORIGINAL REFERENCE NO.: 126:29815a, 29818a

TITLE: Differential response of photosensitized young and old human erythrocytes to photodynamic activation

Rollan, A.; McHale, A. P.

CORPORATE SOURCE: Biotechnology Research Group, School of Applied Biological and Chemical Sciences, University of

Ulster, Coleraine Co. Londonderry, BT52 1SA, UK Cancer Letters (Shannon, Ireland) (1996), Volume Date

1997, 111(1,2), 207-213

CODEN: CALEDQ; ISSN: 0304-3835

Elsevier

PUBLISHER: DOCUMENT TYPE: Journal LANGUAGE: English

AUTHOR(S):

SOURCE:

It has recently been proposed that photosensitized erythrocytes may play an important role in the delivery and targeting of agents such as photosensitizers and chemotherapeutics for use in cancer treatment. It has been suggested that loading of photosensitized erythrocytes with chemotherapeutic agents would provide an ideal means of combining both treatment modalities. The recent application of real-time confocal laser scanning microscopy to the study of immediate effects of photodynamic activation on photosensitized erythrocytes has enabled us, in this study, to distinguish between the differential susceptibility of age-d. resolved sub-populations of human erythrocytes to photodynamic activation. In this study we demonstrate that younger (low age-d.) sub-populations of photosensitized erythrocytes are less susceptible than older (high age-d.) sub-populations to photodynamic activation. We also demonstrate that this phenomenon is exhibited by cells photosensitized using hematoporphyrin derivative and rose bengal as photosensitizers. In both cases no significant difference in uptake of photosensitizer by both populations could be observed using absorbance spectrophotometry. The study suggests that age-d. resolution of erythrocytes prior to loading and photosensitization might provide a means of enhancing the release of loaded components from the photosensitized system and this would, in turn, enhance the potential use of photosensitized erythrocytes as delivery or

targeting systems for use in combination cancer therapies. OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS) REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 15 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1996:467217 CAPLUS

DOCUMENT NUMBER: 125:137244

ORIGINAL REFERENCE NO.: 125:25577a,25580a

TITLE: Gels for encapsulation of biological materials INVENTOR(S): Hubbell, Jeffrey A.; Pathak, Chandrashekhar P.;

Sawhney, Amarpreet S.; Desai, Neil P.; Hossainy, Syed

F. A.

PATENT ASSIGNEE(S): University of Texas System, USA

SOURCE: U.S., 34 pp., Cont.-in-part of U.S. Ser. No. 870, 540. CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 13

PATENT INFORMATION:

DATE APPLICATION NO. DATE PATENT NO. KIND US 5529914 A 19960625 US 1992-958870 19921007 US 53232984 A 19930803 US 1991-740632 19910805 US 5380536 A 19950110 US 1991-740703 19910805 CA 2117584 A1 19930902 CA 1993-2117584 19930301 CA 2117584 C 19980922 UN 1993-US1776 19930301

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W: AU, BB, BG, BR, CA, FI, HU, JP, KP, KR, LK, MG, MN, MW, NO, NZ,
                                             PL, RO, RU, SD, SK, UA
                                 RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                    AII 9337809
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                   AU 683209
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                   EP 627912 A1 19941214 EP 1993-907078 EP 627912 B1 20040512
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EP 627912 B1 20040512
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, TE, IT, LI, LU,
JP 07506961 T 1950803 JP 1993-515100
JP 3011767 B2 20000221
US 5573934 A 19961112 US 1993-24557
BR 9306041 A 19971118 BR 1993-6041
AT 266389 T 20040515 AT 1993-907078
PT 627912 E 20040515 AT 1993-907078
PT 627912 E 20040831 PT 1993-907078
US 5858746 A 1999112 US 1995-467693
US 5858747 A 1998110 US 1995-467693
US 58143743 A 19981201 US 1995-467693
US 5801033 A 19981201 US 1995-467693
US 5801033 A 19981201 US 1995-467693
US 6258870 B1 20010710 US 1995-4676815
US 6632446 B1 20010710 US 1997-783387
US 6231892 B1 20010710 US 1997-783387
US 632446 B1 20010710 US 1998-33871
US 6632446 B1 2001014 US 2006-694836
US 20020058318 A1 2002516
US 20030087985 A1 20030508 US 2001-910663
US 70433819 B2 20051226
US 20040138329 A1 20040156 US 2003-607247
US 20040183190 B2 20061226
US 20040183190 A1 20040107 US 2004-761180
US 20040183190 A1 20040107 US 2004-761180
US 20040195710 A1 20041010 US 2008-743687
US 20040195710 A1 20041007 US 2004-761180
US 20080274201 A1 2008100 US 2008-172063
US 7413781 B2 2008019
US 1991-740632 PA
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US 1990-59880 B2 19901015
US 1991-740632 A3 19910805
US 1991-740703 A2 19910805
US 1992-843485 B2 19920228
US 1992-870540 A2 19920420
US 1992-870540 A2 19920420
US 1993-24657 A1 19930301
US 1993-24657 A1 19930301
US 1993-24657 A3 19940428
US 1994-232054 A3 19940428
US 1994-336393 A3 1994110
US 1995-379848 A2 19950127
US 1995-467693 A3 19950607
US 1995-475175 A2 19950607
US 1995-844160 B3 19950607
US 1995-810089 B1 19950607
US 1995-81089 B1 19950801
US 1997-878387 A1 19970113
US 2000-694836 A1 20001023
US 2001-811901 B2 20010319
US 2001-811663 B1 20010719
US 2001-910663 B1 20010719
US 2001-916663 B1 20010719
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                                                                                                                                                     US 2004-761180
                                                                                                                                                     US 2004-761180 A3 20040120
US 2006-644606 A1 20061222
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
AB This invention provides novel methods for the formation of biocompatible
membranes around biol. materials using photopolymm. of water-soluble mols.
The membranes can be used as a covering to encapsulate biol. materials or
biomedical devices, as a 'glue' to cause >1 biol. substance to adhere
together, or as carriers for biol. active species. Several methods for

forming these membranes are provided. Each of these methods utilizes a polymerization system containing water-soluble macromers, species which are at once

polymers and macromols. capable of further polymerization The macromers are polymerized by using a photoinitiator (such as a dye), optionally a cocatalyst, optionally an accelerator, and radiation in the form of visible or long-wavelength UV light. The reaction occurs either by suspension polymerization or by interfacial polymerization. The polymer membrane can be

formed directly on the surface of the biol, material, or it can be formed on material which is already encapsulated.

OS.CITING REF COUNT: 54 THERE ARE 54 CAPLUS RECORDS THAT CITE THIS

RECORD (62 CITINGS)

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 16 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1996:418715 CAPLUS

DOCUMENT NUMBER: 125:109068

ORIGINAL REFERENCE NO.: 125:20327a,20330a

TITLE: Single cravfish neuron as a new test-object for search

and examination of PDT photosensitizers

AUTHOR(S): Uzdensky, Anatoly B.; Kutko, Olga Yu.; Pasikova,

Natalya V.

Dept. Biophysics and Biocybernetics, Rostov State CORPORATE SOURCE:

University, Rostov-on-Don, 344104, Russia Proceedings of SPIE-The International Society for SOURCE:

Optical Engineering (1996), 2625(Photochemistry:

Photodynamic Therapy and Other Modalities), 512-518 CODEN: PSISDG; ISSN: 0277-786X

PUBLISHER: SPIE-The International Society for Optical Engineering

DOCUMENT TYPE: Journal LANGUAGE: English

An isolated crayfish stretch receptor neuron was used as a new test-object for cytophysiol. study of various photosensitizers. This large cell is very suitable for complex electrophysiol, and cytol, investigation. It generates spikes with a nearly constant frequency, and dynamics of impulse activity shifts under the laser irradiation may be precisely studied at this stable background. The exptl. procedure was as follows: 30 min control spike frequency registration - 30 min neuron staining - He-Ne-laser irradiation with continuous registration of cell response dynamics. typical response of photosensitized neuron to laser irradiation was impulse activity acceleration after some latency and then irreversible block of spike generation. Dependencies of spike frequency acceleration and neuron lifetime on photosensitizer concentration allowed to compare different photosensitizer efficiencies. As the first set of photosensitizers methylene blue, janus green, rose bengal, and chlorin e6, were studied. Chlorin e6 was most potent photosensitizer among them. Such approach provides evaluation of both; initial threshold alteration in cell membrane

and cytotoxic events leading to the cell death. THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD OS.CITING REF COUNT: (5 CITINGS)

L6 ANSWER 17 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

1995:818777 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 123:222385 ORIGINAL REFERENCE NO.: 123:39507a,39510a

TITLE: Agent for visual marking of body tissues

INVENTOR(S): Heywang-Koebrunner, Sylvia; Weitschies, Werner; Speck, Ulrich; Fritzsch, Thomas

PATENT ASSIGNEE(S): Schering A.-G., Germany

SOURCE: Ger. Offen., 5 pp. CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE APPLICATION NO. PATENT NO. ----A1 19950810 DE 1994-4403789 DE 4403789 19940203 A1 19950810 CA 1995-2182686 CA 2182686 WO 9520981 A1 19950810 WO 1995-EP123 W: CA, JP, US RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE A1 19961120 EP 1995-906937 19950113 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE JP 09508397 T 19970826 JP 1995-520342 19950113 PRIORITY APPLN. INFO.: DE 1994-4403789 A 19940203 WO 1995-EP123 W 19950113

AB The invention concerns the use of colored NMR or x-ray contrast media or of dye-containing ultrasound contrast media for the preparation

of diagnostic agents for the visual marking of body tissues. Some possible agents that are discussed are: NMR (metalloporphyrins, iron oxide particles, nitroxides, melanin): x-ray (Rose Bengal.

erythrosin, tetrachlorotetraiodofluorescein); and ultrasound (dye-containing ultrasound contrast media microparticles composed of a covering of a biol. degradable polymer and a gas- and dye-containing center).

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

L6 ANSWER 18 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1995:786246 CAPLUS

DOCUMENT NUMBER: 123:192564

ORIGINAL REFERENCE NO.: 123:34165a,34168a

TITLE: Protective effect of amphotericin B against lethal

photodynamic treatment in yeast
AUTHOR(S): Lazarova, Galina; Tashiro, Hideo

CORPORATE SOURCE: Inst. Microbiol., Bulgarian Acad. Sci., Sofia, 1113,

Bulg.

SOURCE: Microbios (1995), 82(332), 187-96 CODEN: MCBIA7; ISSN: 0026-2633

PUBLISHER: Faculty Press

DOCUMENT TYPE: Journal LANGUAGE: English

The effect of polyenia antibiotic amphotericin B on photodynamically induced cell damage was investigated using Kluyveromyces fragilis. The photosensitizers applied are known to act via cell membrane damage (rose bengal and toluidine blue) or via DNA modification causing genotoxic effects (8-methoxypsoralen). Methylene blue was shown to cause membrane damage comparable with the effect of rose bengal and toluidine blue. Under conditions of photodynamic damage a pronounced protective effect of the antibiotic was evident in increased cell survival with all of the photosensitizers tested. Mitochondrial activity indicated a tendency of the antibiotic to protect the cells. The protective role of amphotericin B is discussed in the light of possible implications for photodynamic therapy of microbial infections.

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD
(6 CITINGS)

L6 ANSWER 19 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1995:494630 CAPLUS

DOCUMENT NUMBER: 122:234390

ORIGINAL REFERENCE NO.: 122:42711a,42714a

TITLE: Photosensitization method of inactivation of viral and

bacterial blood contaminants

INVENTOR(S): Platz, Matthew S.; Goodrich, Raymond P., Jr.; Yerram,

Nagendar

Cryopharm Corp., USA PATENT ASSIGNEE(S):

PCT Int. Appl., 169 pp. SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 12

PATENT INFORMATION:

PA	TENT I	.00			KIN	D	DATE			APPI	LICAT	ION I	NO.		D	ATE	
WO	9502	324			A1		1995	0126		wo :	1994-	US74	99		1	9940	706
	W:	ΑT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,
		JP,	KΡ,	KR,	ΚZ,	LK,	LU,	MG,	MN,	MW,	NL,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SK,	UA,	VN											
	RW:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	ΝE,	SN,	TD,	TG		
US	5418	130			A		1995			US :	1993-	9167	4		1	9930	713
AU	9472	177			A		1995	0213		AU :	1994-	7217	7		1	9940	706
PRIORIT:	APP:	LN.	INFO	.:						US :	1993-	9167	4	I		9930	
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											1990-					9901:	
										US :	1991-	6562	54			99102	
											1991-					9910	
										US :	1992-	8256	91	I	A 1	9920:	127

W 19940706 ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 122:234390

AB A method is provided for inactivating viral and/or bacterial contamination

US 1993-47749

WO 1994-US7499

A 19930414

in blood cellular matter, e.g. erythrocytes, platelets, or protein fractions. The cells or protein fractions are mixed with chemical

sensitizers and irradiated with e.g. UV, visible, gamma, or x-

ray radiation. Preparation of some sensitizer compds. is

included, as are inactivation studies.

OS.CITING REF COUNT: THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(7 CITINGS) REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 20 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

1994:239238 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 120:239238 ORIGINAL REFERENCE NO.: 120:42241a,42244a

TITLE: Photodynamic therapy mediated induction of early

response genes

Luna, Marian C.; Wong, Sam; Gomer, Charles J. AUTHOR(S): Clayton Ocular Oncol. Cent., Child. Hosp., Los

CORPORATE SOURCE: Angeles, CA, 90027, USA

Cancer Research (1994), 54(5), 1374-80 SOURCE:

CODEN: CNREA8: ISSN: 0008-5472

DOCUMENT TYPE: Journal

LANGUAGE: English

Photodynamic therapy (PDT) generates reactive oxygen species which initiate the cytotoxic events of this tumor treatment. The

authors demonstrate that PDT mediated oxidative stress induced a transient increase in the early response genes c-fos, c-jun, c-myc, and erg-1 in murine radiation-induced fibrosarcoma cells. Incubation of

exponentially growing cells with porphyrin based photosensitizers in the dark also induced an increase in the mRNA levels of early response genes. However, the xanthine photosensitizer, rose bengal, produced increased c-fos mRNA levels only following light treatment. Nuclear runoff expts. confirmed that the induction of c-fos mRNA is controlled in part at the level of transcription. Likewise, a chloramphenicol acetyltransferase reporter construct containing the major c-fos transcriptional response elements was inducible by porphyrin and PDT. Signal transduction pathways associated with PDT mediated c-fos activation were examined by treating cells with protein kinase inhibitors. Staurosporine and 1-(5-isoquinolinesulfonyl)-2-methylpiperazine inhibited PDT mediated c-fos activation while N-(2-quanidinoethyl)-5-isoquinoline-sulfonamide had no effect. In addition, quinacrine, which can inhibit phospholipase activity, blocked PDT induced c-fos mRNA expression. These results suggest that photosensitizer mediated oxidative stress acts through protein kinase-mediated signal transduction pathway(s) to activated early response genes.

OS.CITING REF COUNT: 74 THERE ARE 74 CAPLUS RECORDS THAT CITE THIS RECORD (74 CITINGS)

L6 ANSWER 21 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1991:20198 CAPLUS DOCUMENT NUMBER: 114:20198

ORIGINAL REFERENCE NO.: 114:3545a,3548a

TITLE: Primary effects of singlet oxygen sensitizers on eggs

and embryos of sea urchins

AUTHOR(S): Marthy, Hans Juerg; Murasecco-Suardi, Patricia;

Oliveros, Esther; Braun, Andre M.

CORPORATE SOURCE: Lab. Arago, Univ. Pierre et Marie Curie, Banyuls-sur-Mer, 66650, Fr.

SOURCE: Journal of Photochemistry and Photobiology, B:

Biology (1990), 7(2-4), 303-15

CODEN: JPPBEG; ISSN: 1011-1344

DOCUMENT TYPE: Journal LANGUAGE: English

AB Photodynamic effects of soes bengal, a well-known singlet O sensitizer, and of hematoporphyrin derivative, the most widely used sensitizer in photodynamic therapy of tumors, could be visualized using sea urchin eggs and embryos. This biol. material is a valuable model for the anal. of mechanisms and/or sites of the photodynamic action occurring in any living tissue. Depending on the sensitizer used, singlet O may be identified as the main mediator of the cytotoxic effects observed Besides observations made on the living, in particular within the context of fertilization ability of the egg cell, gross damages of the cells are morphol. analyzed by SEM. The results support the working hypothesis explaining the different susceptibility of healthy and tumor

cells for photosensitization as a cell cycle phenomenon.

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD
(3 CITINGS)

L6 ANSWER 22 OF 22 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1988:461480 CAPLUS DOCUMENT NUMBER: 109:61480

ORIGINAL REFERENCE NO.: 109:10213a,10216a

TITLE: Increase of marking stability of radionuclide-marked carrier materials

INVENTOR(S): Wunderlich, Gerd; Dreyer, Rolf; Fischer, Steffen;
Beyer, Renate

PATENT ASSIGNEE(S): Medizinische Akademie "Carl Gustav Carus", Ger. Dem.

Rep.

SOURCE: Ger. (East), 3 pp.

CODEN: GEXXA8

DOCUMENT TYPE: Patent LANGUAGE: German FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DD 251745 A1 19871125 DD 1986-289719 19860429
PRIORITY APPLN. INFO.: DD 1986-289719 19860429

AB Radioactive particles permit the internal radiation of surrounded space and inoperable tumors. Radionuclide-marked carrier materials are treated with dissolved organic substances, whereby the adhesion of the radionuclide on the carrier is increased. Human serum albumin after marking with a radionuclide such as I-125, I-131, or At-211 was incubated in 1% aqueous Titan yellow, bromphenol blue, bengal rose, or Alizarin S with agitation at room temperature The process was repeated with another organic substance from those listed above. Centrifuged treated protein particles were washed with distilled H20 and physiol. NaCl solution After suspension of the treated microspheres in physiol. NaCl solution, the preparation was ready to be injected.

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(FILE 'HOME' ENTERED AT 12:30:32 ON 03 MAR 2011)

FILE 'REGISTRY' ENTERED AT 12:30:46 ON 03 MAR 2011 1 S ROSE BENGAL/CN

FILE 'CAPLUS' ENTERED AT 12:31:06 ON 03 MAR 2011

L2 3080 S L1

90 S L2 AND (CANCER OR TUMOR OR TUMOUR OR NEOPLASM)

L4 22 S L3 AND (RADIATION OR RADIOTHERAPY OR X-RAY OR IRRADIATION)

L5 22 DUP REM L4 (0 DUPLICATES REMOVED)

L6 22 S L5

L7 1 S L5 AND (MONOCHROMATIC OR AUGER)

=>

L3

---Logging off of STN---

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Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS

FULL ESTIMATED COST

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

CA SUBSCRIBER PRICE

SUBSCRIBER PRICE

SUBSCRIBER PRICE

SINCE FILE
ENTRY
SESSION
7-20.01
7-20.01
7-20.01
7-20.01

STN INTERNATIONAL LOGOFF AT 12:32:47 ON 03 MAR 2011